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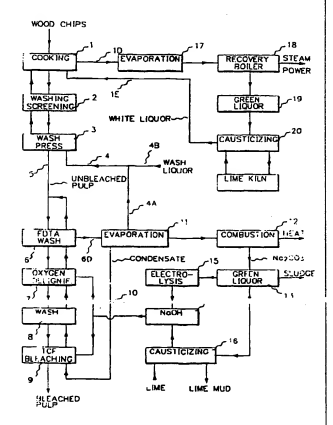
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(54) Title: METHOD OF BLEACHING PULP WITHOUT USING CHLORINE-CONTAINING CHEMICALS

(57) Abstract

The present invention relates to a process for producing pulp which is cooked under alkaline conditions and which is bleached without using chlorine-containing bleaching chemicals, in which process used cooking chemicals are recovered in a first recovery installation (18) and used bleaching chemicals are recovered in a second recovery installation (12), and, preferably, these bleaching chemicals are regenerated, wholly or in part, and reused in an oxygen-delignification stage (7) and/or in a TCF bleaching stage (9).



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WO 94/12720 PCT/SE93/00925

Method of bleaching pulp without using chlorinecontaining chemicals

5 Technical field

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The present invention relates to a process for producing pulp which is cooked under alkaline conditions and which is bleached without chlorine-containing bleaching chemicals, with separate installations being used for recovering and for incinerating/gasifying filtrates from the cooking and the bleaching, respectively.

15 <u>Background and problems</u>

Bleaching pulp without using chlorinecontaining chemicals provides processing solutions which are of great interest from the environmental point of view. Thus, the discharge of impurities into bleaching plant effluent can be reduced substantially and, in the optimum case, virtually eliminated by, in an appropriate manner, collecting up and concentrating the organic and inorganic compounds in the effluent and subsequently incinerating together the concentrates and the used chemicals from the cooking procedure. A process of this type has been reported in Swedish Patent Application no. 9201477-8.

Bleaching without chlorine-containing chemicals is carried out using hydrogen peroxide, sodium hydroxide and ozone, in particular. These bleaching chemicals are expensive, so that bleaching costs are significantly higher than in the case of conventional bleaching using chlorine and chlorine dioxide. It is therefore desirable, if possible, to be able to recover the bleaching chemicals, especially sodium hydroxide, separately. Sodium hydroxide is principally obtained in association with preparing chlorine gas by the

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electrolysis of sodium chloride. With decreasing use of chlorine gas, there is some risk of sodium hydroxide becoming an item in short supply.

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The process, which is advantageous from the environmental point of view, of returning the used bleaching chemicals, and organic material released during the bleaching, to the system for recovering cooking chemicals can give rise to problems under certain circumstances. If the quantity of sodium compounds and sulphur-containing compounds in bleaching plant effluent exceeds the requirement for chemicals to cover losses, "make-up" an arises in the Na/S ratio in the recovery cycle for cooking chemicals. This can lead to problems of having too high an emission of sulphur to the environment, and to other disturbances of the process as well. Another problem, which is equally serious in many works, can be that organic material in the bleaching plant effluent, which is incinerated in the works recovery boiler, leads to overloading of the boiler. A works recovery maximum is often utilized to Overloading therefore results in the production of pulp having to be decreased, which is an economic disadvaninstances where transfer of those In bleaching chemicals to the recovery cycle for cooking chemicals can lead to imbalance in the Na/S ratio and/or overloading of an existing recovery boiler, a separate recovery cycle for the bleaching chemicals would be highly advantageous.

Solution and advantages

In accordance with the process which is indicated in Patent Claim 1, this invention provides a solution, which is technically and economically advantageous, both to the requirement for recovering bleaching chemicals separately and to the problems of limited capacity in the recovery boiler. In the



following, the term bleaching also embraces oxygen delignification.

Brief description of the figures

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The invention will be described below with reference to the enclosed figures, in which:

Fig. 1 shows a block diagram of a preferred embodiment in principle according to the invention, and Fig. 2 shows a preferred installation for its realization.

Detailed description

Figure 1 shows an embodiment of the invention in the form of a block diagram of a sulphate works in which bleaching is totally chlorine-free the (henceforth termed TCF). Wood chippings are cooked in a house 1 which is equipped for sulphate cooking and which permits delignification to low kappa numbers. Subsequently, the pulp is washed and screened 2. After the screening 2, there follows a further washing stage 3. A preferred form of that invention the final washing for is stage unbleached pulp consists of a washing press 3. another press, which permits dehydration to a high dry matter content. The washing liquid 4 for the final washing stage 3 consists of chemically purified water 4B and/or evaporation condensate 4A. The unbleached pulp 5 is thus thoroughly washed.

After the washing, and any intermediate storage, the pulp is pumped to an acid wash stage 6, which includes the possible addition of a chelating agent, for example EDTA. The purpose of this stage is to remove heavy metals. Filtrate from a suitable stage in the TCF bleaching plant 9 and, where appropriate, recirculated filtrate from the wash 6 after the acid wash stage 6 are used for adjusting the consistency of

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the pulp. After the acidification, the pulp is washed in a washing system possessing a high degree of washing efficacy. The washing liquid for the wash of the acid wash stage consists of a suitable filtrate from the subsequent oxygen-delignification stage 7 or the TCF bleaching plant 9.

After the acid wash stage, there follows an oxygen-delignification stage 7. In the oxygen stage, delignification takes place to a low kappa number which, for soft wood, is lower than 15, preferably lower than 12, and for hard wood is lower than 12, preferably lower than 10.

The treatment of the pulp with acid and, where appropriate, with chelating agent prior to the oxygen stage 7 removes heavy metal ions which otherwise impair selectivity during the oxygen delignification, i.e. cause degradation of the cellulose. This wash thus permits the delignification to be taken further, which is a major advantage for subsequent TCF bleaching.

After the oxygen-delignification stage, the pulp is washed and then conveyed to the TCF bleaching plant 9. The latter can be designed with a number of different sequences, for example one or more peroxide stages (P), peroxide followed by ozone (PZ), peroxide, ozone and peroxide (PZP), or Z(EOP)P, i.e. ozone, alkali extraction in the presence of oxygen and, where appropriate, peroxide, and subsequently a peroxide stage.

Heavy metals, which are harmful for both the peroxide stage and the ozone stage, have already been removed in the acid wash stage 6 prior to the oxygen stage 7. It is not therefore necessary, as is done in conventional peroxide bleaching, for example in the Lignox process (SE-B-466061), to treat the pulp with chelating agent immediately prior to the peroxide stage.

Washing liquid, preferably in the form of evaporation condensate 10, is supplied to one of the

stages in the TCF bleaching plant. Except for the filtrate from the ozone stage, the filtrates from the different stages are in the main conveyed in a counter current fashion to the pulp and drawn off via the wash after the acid wash stage 6. This filtrate contains organic material released from the pulp both in the oxygen stage 7 and in the TCF bleaching plant 9, and used bleaching chemicals, principally sodium compounds arising from added sodium hydroxide and sulphur compounds as well, for example derived from sulphuric acid used for pH adjustment. In addition, the filtrate will contain washing losses from the final washing stage for unbleached pulp in the organic material, sodium compounds, sulphides relatively small quantities of heavy metals, which are derived originally from the wood and possibly bound in complex form.

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The filtrate from the acid wash stage 6 conveyed to an evaporation stage 11, where it evaporated to high a dry matter content. evaporation 11 can be carried out in a conventional manner in a multi-step evaporation system with steam heating or by so-called mechanical steam compression, by combinations of mechanical steam compression and evaporation with steam, or by evaporation with the aid of low-grade waste heat in the form of hot water or another heat source. Preferably, the same heat-source (e.g. steam) is used for evaporation of the filtrate from the bleaching as for the spent liquor, whereby preferably the filtrate from the bleaching is preevaporated in one or two units in the end (seen from the steam flow) of the line of evaporation units and the spent liquor is pre-evaporated in one or two inbetween units and preferably thereafter joining the two liquid flows for common evaporation (one, two or three units) in the beginning of the line. Alternatively, the two pre-evaporated liquids are finally evaporated in separate units, whereby at least the remaining heat

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(e.g. steam) from the final evaporation of the spent liquor is used for the pre-evaporation and preferably also the remaining heat from the final evaporation of the bleach-effluent is used by joining it with the heat from the final evaporation of the spent liquor. The condensate 4A, 10, which is obtained during the evaporation, can be utilized as a washing liquid in the TCF bleaching plant and/or for washing the unbleached pulp.

The concentrated filtrate containing organic material and sodium compounds, and possibly sulphur compounds as well, is conveyed to a incineration oven/gasification reactor 12, where the organic material is completely or partially oxidized and the sodium compounds are converted in the main to sodium carbonate and, in the appropriate case, sodium sulphate or sodium sulphide. The oxidation can either be carried out in the presence of an excess of oxygen or in the presence of a deficiency of oxygen, in which latter case the oxidation takes place under reducing conditions. The latter method, which is more preferably carried out in a CHEMREC® reactor, involves gasification of the organic material to form a mixture of carbon monoxide, hydrogen gas and carbon dioxide. The incineration or gasification temperature must be sufficiently high to convert essentially all the carbon gaseous products and sodium carbonate. reducing conditions, sodium sulphide is formed from the sulphur compounds which are supplied.

The sodium salts which are formed during incineration are removed in the form of a smelt, which is dissolved in water and forms so-called green liquor 13. During gasification, an aqueous solution of these salts (green liquor) is obtained by direct liquid cooling of the incineration gases, preferably in a so-called quench system. The incineration heat can be utilized for generating steam and/or preparing hot water. The combustible gases, principally hydrogen gas,

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carbon monoxide and methane, which are formed during the gasification represent a flexible energy source. The carbon monoxide gas which is formed can, by means of the so-called shift reaction, be used for generating further hydrogen gas. The crude hydrogen gas obtained in this way can be purified and used together with oxygen gas for the local preparation of hydrogen peroxide.

The green liquor 13 is filtered carefully in order to separate off precipitated substances which are contrary to the process, such as heavy metals and other impurities. It is then utilized in whole or in part for preparing sodium hydroxide 14. This can be carried out electrolysis 15 or either by by causticization 16. In the latter case, slaked lime is added, which reacts with the sodium carbonate to give sparingly soluble calcium carbonate and The calcium carbonate (lime separated off by filtering and conveyed to a lime kiln for re incineration.

Sodium hydroxide, obtained either by the electrolysis of green liquor or alternatively by caustiis used as a bleaching chemical oxygen-delignification stage 7 and/or for regulating the pH in a peroxide stage (P) or an alkali extraction stage (E) in the TCF bleaching plant 9. The system thus becomes to a large extent self-supporting with regard economically sodium hydroxide, which. is addition, is advantageous. In the system almost entirely closed, with the discharges of organic material and other impurities in the effluent water having in the main been eliminated.

Oxygen, which can also be prepared within the works, is required in addition to the above mentioned chemicals.

The upper part of the block diagram in Figure 1 shows how the spent liquor (the filtrate from the cooking) is conveyed to an evaporation installation 17 in

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which the spent liquor is evaporated, in order, subsequently, to be incinerated in a recovery boiler 18. The heat which is produced in the recovery boiler is conserved in the form of steam, whose energy content is subsequently converted, for example into electric the recovery boiler In 18, the inorganic products contained in the spent liquor form a smelt consisting, in the main, of sodium carbonate and sodium sulphide which are collected from the recovery boiler and dissolved in water in order to form green liquor green liquor is then converted causticization installation 20 such that new cooking liquid, i.e. white liquor (NaOH + Na2S), is formed which is returned to the digester 1.

Sodium sulphate is obtained during the suprastoichiometric incineration of bleaching effluents which contain sulphur-containing compounds. The green liquor produced under these circumstances will therefore contain both sodium carbonate and sodium sulphate. Causticization of the green liquor results in the formation of calcium sulphate, which, however, is relatively soluble as compared with the calcium carbonate (lime sludge). The sodium hydroxide solution will therefore contain a relatively high content of sulphate ions. In addition, the lime consumption will increase and the lime sludge will contain a relatively high content of calcium sulphate, which can interfere with the course of the lime cycle. These difficulties can be avoided by using an organic acid, for example oxalic acid or acetic acid, for adjusting the pH in the acid bleaching stages. The organic acid is incinerated the boiler and does not interfere with causticization reaction. Organic acids can, naturally, also be used in systems which are based on gasification of concentrates of bleaching effluents, and in this way the problems with sulphur can be eliminated completely.

When a green liquor containing both sodium

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carbonate and sodium sulphate is electrolyzed, sulphuric acid and sodium hydroxide are obtained. These chemicals can be separated by means of a suitable membrane and subsequently recirculated independently to the bleaching plant.

Sulphur compounds contained in the mother liquor give rise, in the main, to sodium sulphide, rather than sodium sulphate, when partial incineration under reducing conditions, i.e. gasification, place. However, an aqueous solution of sodium sulphide can be oxidized with oxygen, under pressure and at elevated temperature, sodium sulphate, to and the mixture with sodium carbonate subsequently subjected to electrolysis in accordance with the above description.

Any necessary "make-up" to compensate losses of chemicals in the recovery cycle of bleaching plant can be supplied in the form of fresh NaOH and, where appropriate, sulphuric acid, H2SO4. Chemical surpluses and any necessary regulation of the enriched concentration of substances which are contrary to the process and which derive originally from the wood can take place either by transferring a part of green liquor or residual solution from electrolysis to the system for recovering chemicals for regulating the pH of the effluent water of the works. In the first-mentioned case, most of the impurities, for example heavy metals, are removed by way of the green liquor sludge which is obtained on filtering the green liquor, and then dealt with in a manner which is advantageous from the environmental point of view.

Figure 2 shows an embodiment of the invention in the form of a system solution for the fibre line. Soft wood chippings are steamed 1A and preimpregnated 1B and then cooked in a KAMYR® digester 1C equipped for modified sulphate cooking, which permits delignification to low kappa numbers, 18-22, while preserving strength properties. The pulp, which has

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been partially washed in this process, is washed further firstly in a diffuser 2A and after screening 2B on a filter 3A which functions as a combined thickener and washing apparatus. Subsequently, the pulp is washed and thickened on a washing press 3B to the pulp consistency of 25-35%. Chemically purified water 4B or evaporation condensate 4A is used as the washing The washing liquid is conveyed in counter current to the digester, where it moves upwards, again in counter current, to the withdrawal screen, where is drawn off and conducted liquor 1D evaporation 17 and thence to incineration in a recovery boiler 18 (see Figure 1). The smelt from the boiler is dissolved in water to form green liquor containing mainly sodium carbonate and sodium sulphide. The green liquor is subsequently causticized in a known manner to form white liquor containing mainly sodium hydroxide and sodium sulphide. The white liquor 1E is conveyed to the digester for decomposing the wood chippings.

The pulp leaves the washing press with a consistency of 25-35%. It is well washed and contains only about 3 kg of Na+/ton of pulp and about 10 kg of dissolved organic material. In addition, the pulp contains relatively small quantities of heavy metals.

The pulp from the washing press 3B is then diluted to a consistency of about 10% with filtrate, in the proportion of about 6 tons/ton of pulp, from the ozone stage (Z) wash in the subsequent TCF bleaching plant 9. Using a KAMYR[®] MC pump 6A, acid, for example organic acid or sulphuric acid, is added together with a chelating agent such that a pH of 5-6 is achieved. The temperature in the EDTA stage 6B or Q stage should be 50-90°C and the dwell time 30-60 minutes.

The pulp treated in this way is subsequently washed in a washing apparatus 6C (included in the so-called acid washing stage 6) possessing a high degree of washing efficacy, at least 80%, preferably 90-95%, measured as its effect in separating off manganese. In

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the preferred case, a KAMYR® two-stage diffuser 6C is employed. Other washing devices, for example a washing press or one or more washing filters in series, may be employed. During the washing, the pulp is freed from heavy metals, which are to be found in the form of complexes in the filtrate 6D, which is conveyed to separate evaporation 11, (see Fig. 1) oxidation and chemical recovery. The filtrate 6D also contains most of the organic material which was released in the oxygen-delignification stage 7 and the TCF bleaching 9. In addition, most of the sodium compounds, and, where appropriate, the sulphur compounds which were added to the bleaching are to be found in the filtrate.

NaOH, at the rate of 10-20 kg per ton of pulp, and, where appropriate, magnesium salt, are supplied in an MC pump 7a to the pulp, which is well washed and freed from heavy metals, and the pulp is then conveyed under pressure via an MC mixer 7B, in which oxygen and, where appropriate, steam are supplied, to a reactor 7C having a dwell time of about 30-90 minutes, preferably of about 60 minutes. The temperature of the pulp is 80-110°C and the pressure 3-10 bar. During the oxygen delignification, the kappa number is lowered to less preferably less than 15, 12. The residual chemicals and released organic material are washed out after the oxygen stage in one or more washing devices 8 washing efficacy of having a degree of 80-95%, preferably 90-95%. Figure 2 shows a KAMYR® two-stage diffuser 8, but the washing can also be carried out using other devices having a similar degree of washing efficacy, for example filters or washing presses.

Hydrogen peroxide, at the rate of 10-35 kg of H₂O₂/ton of pulp, preferably 20-30 kg of H₂O₂/ton of pulp and sodium hydroxide, at the rate of 5-30 kg of NaOH/ton of pulp, preferably 15-25 kg of NaOH/ton of pulp, are subsequently supplied to the oxygendelignified pulp by means of an MC pump 9A and the pulp is then heated to a temperature of 75-95°C, preferably

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of 80-90°C. After that, the pulp is conveyed to one or more reaction towers having a dwell time of 3-8 hours, preferably 4-6 hours.

The peroxide-bleached pulp, having a brightness of 75-85 ISO, is washed in a washing apparatus 9C possessing a high degree of washing efficacy, for example in a KAMYR® two-stage diffuser. Either chemically purified water or evaporation condensate 10 is used as the washing liquid. The temperature of the washing liquid should be 35-55°C, preferably 40-50°C. The displaced peroxide-bleaching filtrate 9D is recirculated to the preceding washing apparatus 8, it being possible to utilize both residual peroxide and heat in the peroxide stage.

After the peroxide bleaching stage 9B, the pulp is pumped onwards by means of a KAMYR® MC pump 9E and is acidified to pH 2-6, preferably pH 3-4. Sulphuric acid, for example, or an organic acid, for example acid, acid acetic is used or acidification. Ozone gas in oxygen, at a concentration of 5-15% O3, is added at a pressure of 5-12 bar and mixed into the pulp suspension, which has a consistency of about 10%. One or more mixers of the KAMYR® MC mixer 9F type, or another type of efficient mixing apparatus, is/are employed for the admixture. The temperature of the pulp suspension should be 35-55°C, preferably 40-50°C. The charge of ozone gas should be 2-6 kg of 03/ton of pulp, preferably 3-5 kg of 03/ton of pulp.

After the admixture, the pulp suspension containing ozone gas is conveyed through a reactor 9G having a dwell time of 1-10 minutes, preferably 1-4 minutes. Subsequently, the pressure is lowered in a cyclone device 9H, with the gas being separated from the pulp suspension. The residual gas, which mainly consists of oxygen with small quantities of unreacted ozone, is cleaned of fibre in a scrubber (not shown) and conveyed to an ozone-destroying apparatus. The oxygen gas can be compressed in a compressor and reused

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in the oxygen-delignification stage, for example.

The pulp suspension, which has been freed of gas, is pumped to a washing stage 9i, for example a KAMYR[®] one-stage diffuser. If appropriate, hydroxide, for neutralizing to pH5-10, and sulphur dioxide, for eliminating remaining ozone in the pulp suspension, are added before the wash. After washing stage, alkali and hydrogen peroxide supplied in a $KAMYR^{\textcircled{R}}$ MC pump and/or mixer 9J. charge of peroxide should correspond to 1-5 kg of H2O2 per ton of pulp and the charge of alkali should be sufficiently high to adjust the pH to pH 10-11. The temperature of the peroxide stage 9K must be 50-80°C, preferably 60-75°C, and the dwell time of the pulp 1-4 hours, preferably 2-3 hours.

After the peroxide stage 9K, the pulp is washed in a KAMYR[®] one-stage diffuser 9L or other washing device having a similar degree of washing efficacy. At this stage, the brightness of the finally bleached pulp is 85-90 ISO, preferably 88-90 ISO.

If sulphuric acid is used in the acid wash stage 6 and the ozone stage 9B, the filtrate 6D, which is withdrawn from the acid wash stage (the Q stage), will have the following approximate composition per ton of pulp:

Organic material	about 75 kg
Na+	" 30 kg
SO ₄	<u>" 15 kg</u>
Total dry matter	120 kg

The quantity of liquid is about 10 m³ per ton of pulp. This corresponds to a dry matter content of about 1.2%. The filtrate is evaporated to a dry matter content of 50-70%, with about 9 tons of condensate being obtained which is used for washing the pulp after the first peroxide stage, or at another suitable point in the process. The evaporated concentrate, which has a calorific value of about 9 MJ/kg of dry matter, is incinerated in an oxidizing environment, a smelt

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containing about 22 kg of Na2SO4 and about 52 kg of Na₂CO₃ being formed. The smelt is dissolved in water. The "green liquor" which is obtained is thoroughly filtered in order to separate off solid impurities, for example by precipitating out heavy metal salts. electrolyzed is solution Subsequently, the electrolyzers having membranes which separate sodium hydroxide and sulphuric acid which are formed. Based on the chemical content of the green liquor, about 13 kg of sulphuric acid and about 46 kg of NaOH are thus formed at an efficiency of 90%. The necessary requirement for the oxygen-delignification stage and the TCF bleaching is about 15 kg of sulphuric acid per ton of pulp and about 50 kg of NaOH per ton of pulp. requirement for fresh chemicals is therefore limited to about 2 kg of H2SO4 and 4 kg of NaOH, in addition to peroxide and oxygen gas.

If an organic acid is used for acidification in the acid wash stage and ozone stage, it is in the main only sodium carbonate which is obtained in the smelt. After dissolution in water, the sodium carbonate solution can be treated with slaked lime, with sodium hydroxide and calcium carbonate (lime sludge) being formed. The latter is washed and transferred to the lime kiln for re-incineration. The sodium hydroxide is used for bleaching.

motivation for there is no economic If recovering sodium hydroxide or sulphuric acid, the salts can be used for other purposes, for example as make-up chemicals in the cooking chemical cycle. this case, the benefit is obtained that the recovery boiler for the cooking chemical cycle is not loaded the organic material released in oxygenwith delignification stage and/or in the TCF bleaching plant. In addition, discharge of organic material from the bleaching plant is eliminated.

Naturally, the invention can also be used in connection with the production of pulp from hard wood

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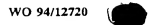
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or other raw material, such as annual plants. requirement for chemicals will vary depending on the pulp brightness required and the on cellulose-containing raw material which is incorporated. The method can also be used in those cases where the cooking liquid is sulphur-free or has a low sulphur content and is mainly made up, for example, of an alkaline hydroxide. The method can advantageously be employed in those cases where the alkali metal base in the cooking and/or the oxygen delignification and TCF bleaching mainly consists of potassium instead of sodium. It will be evident to the person skilled in the appurtenant incineration/gasification art that the installation 12 can operate in accordance essentially any of the currently incinknown eration/gasification principles, a CHEMREC® even if reactor is preferred. In addition, it will be evident that the recovery installation intended for the spent liquor can consist of a gasification reactor, for example a CHEMREC® reactor. Over and above this, should be pointed out that the invention is not limited to only two recovery boilers/reactors and that it is possible to conceive of combining certain parts in the recovery cycle from the systems illustrated separately in Figure 1. The degree of washing efficacy is defined as follows:

(X-Y) @ 100; where X is the quantity of unwanted X substance prior to washing and Y is the quantity of the said substance remaining after washing, for a given quantity of pulp. The manganese content is advantageously employed as a reference value for the said substance.

The solitary "A" in Figure 2 indicates the addition of chemically purified water or evaporation condensate.



Patent Claims

- Process for producing pulp which is cooked under alkaline conditions and which in the main is bleached without using chlorine-containing bleaching chemicals, c h a r a c t e r i z e d i n that used cooking chemicals are recovered in a first recovery installation (18) used bleaching chemicals are recovered, wholly or in part, in a second recovery installation (12), which bleaching chemicals are preferably regenerated, wholly or in part, and reused in an oxygen-delignification
- 15 2. Process according to Patent Claim 1, c h a r a c t e r i z e d i n that prior to an oxygen-delignification stage (7), the unbleached pulp is treated with acid and, where appropriate, with chelating agent and is washed (6), preferably with a 20 degree of washing efficacy of at least 80%, more preferably at least 90%.

stage (7) and/or in a TCF bleaching stage (9).

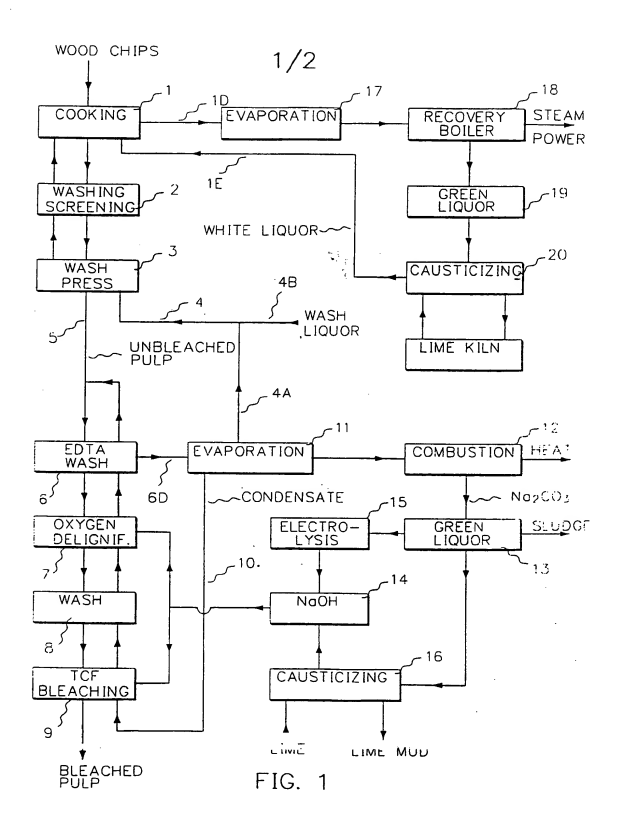
- 3. Process according to Patent Claim 2, c h a r a c t e r i z e d i n that the greater part of the organic material released in the oxygen-delignification stage (7) and in the TCF bleaching stage (9), and of the bleaching chemicals, which have been supplied and in the main used, is drawn off together with the filtrate (6D) from the said washing stage (6C).
- 4. Process according to Patent Claim 3, characterized in that the filtrate (6D) which is drawn off is limited to at most 15 tons per ton of pulp, preferably at most 10 tons per ton of pulp, and preferably the content of dry matter in the drawn-off filtrate is concentrated by evaporation to at least 50%, preferably at least 60%.

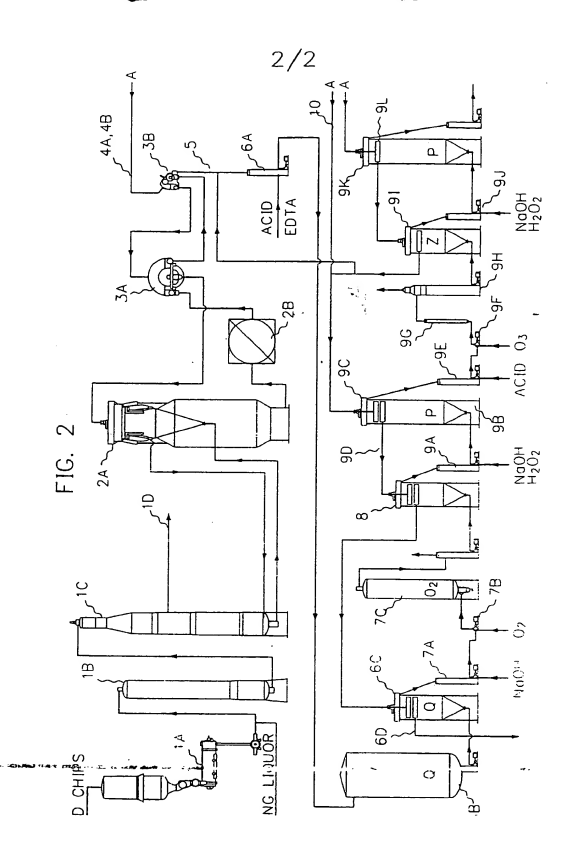
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- 5. Process according to Patent Claim 4, c h a r a c t e r i z e d i n that the concentrated filtrate (6D) which is drawn of from the bleaching is incinerated in an oxidizing environment under stoichiometric or suprastoichiometric conditions, and the resulting inorganic chemicals are dissolved in liquid.
- 6. Process according to Patent Claim 4,
 10 c h a r a c t e r i z e d i n that the filtrate (6D)
 which is drawn off from the bleaching and concentrated
 is gasified in a reducing environment, most preferably
 in a CHEMREC® reactor, and the resulting imorganic
 chemicals are dissolved in liquid.
 - 7. Process according to Patent Claim 5 or 6, c h a r a c t e r i z e d i n that alkali metal hydroxide is recovered from the liquid solution by causticization or by electrolysis of dissolved alkali metal carbonate.
- 8. Process according to Patent Claim 5 or 6, c h a r a c t e r i z e d i n that sulphuric acid is recovered from the liquid solution by electrolysis of dissolved alkali metal sulphate.
 - 9. Process according to Patent Claim 6, c h a r a c t e r i z e d i n that dissolved sulphide obtained by reductive incineration is oxidized with oxygen to sulphate prior to electrolysis.
 - 10. Process according to Patent Claims 1 to 7, c h a r a c t e r i z e d i n that necessary acidification in the acid wash stage (6), with or without the addition of chelating agent, and any acid stages in the TCF bleaching plant (9), are brought about by supplying an organic acid.

11. Process according to Patent Claim 1, c h a r a c t e r i z e d i n that, the bleach effluent and the spent liquor are evaporated by using mainly the same heat source.





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A. CLASSI	FICATION OF SUBJECT MATTER					
IPC5: D2	1C 9/00, D21C 11/00 International Patent Classification (IPC) or to both	national classification and IPC				
B. FIELDS	SEARCHED					
Minimum doc	umentation searched (classification system followed	by classification symbols)				
IPC5: D2:	1C					
Documentation	n searched other than minimum documentation to t	he extent that such documents are included i	n the fields searched			
SE,DK,FI	,NO classes as above					
Electronic data	a base consulted during the international search (nar	ne of data base and, where practicable, search	h terms used)			
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	•				
Category* C	Itation of document, with indication, where a	ppropriate, of the relevant passages -	Relevant to claim No.			
A L	JS, A, 5164043 (BRUCE F. GRIGGS 17 November 1992 (17.11.92)	ET AL),	1			
A U	US, A, 5164044 (BRUCE F. GRIGGS	ET AL),	1			
	17 November 1992 (17.11.92)					
	4444000					
Further documents are listed in the continuation of Box C. X See patent family annex.						
Special categories of cited documents To later document published after the international filling date or pri						
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INITATIONAL SEARCH REPORT



International application No. PCT/SE 93/00925

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